1 Improved Prediction of Dimethyl Sulfide (DMS) Distributions in the

2 NE Subarctic Pacific using Machine Learning Algorithms

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Abstract. Dimethyl sulfide (DMS) is a volatile biogenic gas with the potential to influence regional climate as a 8 9 source of atmospheric aerosols and cloud condensation nuclei (CCN). The complexity of the oceanic DMS cycle 10 presents a challenge in accurately predicting sea-surface concentrations and sea-air fluxes of this gas. In this study, 11 we applied machine learning methods to model the distribution of DMS in the NE Subarctic Pacific (NESAP), a 12 global DMS hot-spot. Using nearly two decades of ship-based DMS observations, combined with satellite-derived 13 oceanographic data, we constructed ensembles of 1000 machine-learning models using two techniques, random forest regression (RFR) and artificial neural networks (ANN). Our models dramatically improve upon existing 14 15 statistical DMS models, capturing up to 62% of observed DMS variability in the NESAP and demonstrating notable 16 regional patterns that are associated with mesoscale oceanographic variability. In particular, our results indicate a 17 strong coherence between DMS concentrations, sea surface nitrate (SSN) concentrations, photosynthetically active 18 radiation (PAR) and sea surface height anomalies (SSHA), suggesting that NESAP DMS cycling is primarily influenced by heterogenous nutrient availability, light-dependent processes and physical mixing. Based on our 19 20 model output, we derive summertime, sea-air flux estimates of 1.16±1.22 Tg S in the NESAP. Our work 21 demonstrates a new approach to capturing spatial and temporal patterns in DMS variability, which is likely 22 applicable to other oceanic regions.

23 1 Introduction

24 Dimethyl sulfide (DMS), a volatile biogenic gas, is an important component of the marine sulfur cycle. 25 This molecule is an important substrate for specific methylotrophic bacteria (Vila-Costa et al., 2006; Lidbury et al., 26 2016; Green et al., 2011; Hatton et al., 2012), with a recognized importance to marine microbial metabolism (Vila-27 Costa et al., 2006) and food web interactions (Nevitt, 2008). Moreover, DMS constitutes the largest fraction of 28 bulk non-sea salt (NSS) sulfate emissions to the atmosphere (Bates et al., 1992; Ksionzek et al., 2016), where it is 29 rapidly oxidized to form aerosols that act as cloud condensation nuclei (CCN: Charlson et al., 1987; Hegg et al., 30 1991; Korhonen et al., 2008), potentially influencing regional albedo and climate (Charlson et al., 1987; Ayers and 31 Cainey, 2007). Given the ecological roles of DMS and its potential influence on global climate, substantial research 32 has focused on characterizing the dynamics of this compound in seawater. This work has revealed considerable 33 complexity in the oceanic DMS cycle, which has limited the development of simple predictive algorithms 34 describing its spatial and temporal variability.

Oceanic DMS production and loss are tightly linked with the biological cycling of the related metabolites
 dimethyl sulfoniopropionate (DMSP) and dimethyl sulfoxide (DMSO). DMS is believed to be primarily derived

37 from the cleavage of DMSP (Kiene and Linn, 2000), but it can also be cycled through biological DMSO reduction (Spiese et al., 2009) and oxidation (Lidbury et al., 2016), and abiotically by light-dependent reactions (del Valle et 38 39 al., 2007; Royer et al., 2016). DMS cycling is influenced by a suite of environmental and ecological factors, including release from phytoplankton cells into the dissolved pool via grazing (Dacev and Wakeham, 1986), viral 40 lysis (Malin et al., 1998), or exudation. Oxidative stress generated by other variables such as temperature (Kirst et 41 42 al., 1991), salinity (Dickson and Kirst, 1987), UV radiation (Kinsey et al., 2016), and nutrient limitation (Bucciarelli 43 et al., 2013; Spiese & Tatarkov, 2014) may also enhance the cycling of DMSP and DMSO, which may regulate 44 DMS concentrations through cascading oxidative pathways (Sunda et al., 2002). Finally, variability in surface wind 45 fields can modulate the rates of DMS sea-air exchange, providing a significant source of heterogeneity in surface 46 water DMS concentrations (Rover et al., 2016). These examples illustrate the complex non-linearity of the oceanic 47 DMS cycle.

48 Over the past two decades, a number of approaches have been developed to model DMS distributions at 49 both global (Bock et al., 2021; Galí et al., 2018; Simó and Dachs, 2002; Vallina and Simó, 2007) and regional 50 (Watanabe et al., 2007) scales. These models have been largely based on linear regression techniques estimating 51 DMS concentrations using one or two predictors. To date, these studies have focused on a number of variables, 52 including ratio of chlorophyll a (Chl-a) to mixed layer depth (MLD) (Simó and Dachs, 2002), sea surface 53 temperature (SST) and nitrate (SSN) (Watanabe et al., 2007), solar radiation dose (SRD) (Vallina and Simó, 2007), 54 photosynthetically active radiation (PAR) and modelled DMSP concentrations (Galí et al., 2018). Some of these 55 models have demonstrated reasonably good performance at global scales, but their predictive power is generally 56 diminished at regional scales (Herr et al., 2019), failing to accurately resolve important smaller-scale features 57 (Belviso et al., 2003; Nemcek et al., 2008; Royer et al., 2015; Tortell, 2005b).

58 In recent years, machine-learning algorithms have been increasingly used to derive predictions for non-59 linear oceanic systems. For example, these methods have been successfully applied to describe the spatial and 60 temporal patterns of global methane flux (Weber et al., 2019), nitrous oxide dynamics (Yang et al., 2020), and 61 carbon export (Roshan and DeVries, 2017). To our knowledge, only two studies have thus far applied machine-62 learning to describe DMS distributions, with one study focused on the Arctic (Humphries et al., 2012) and the other 63 exploring a global domain (Wang et al., 2020). Despite producing algorithms with reasonable predictive skill, these 64 two studies found limited success in resolving the underlying relationships driving DMS variability. This was 65 partially due to a reliance on indirect sensitivity tests assessing the importance of predictor variables, and also, 66 potentially, from the large-scale averaging applied to the underlying data fields (1x1°; 111 km²). Analyses at higher spatial resolution may reveal mesoscale (roughly 20-200 km) and sub-mesoscale (roughly 1-20 km) patterns that
would otherwise be obscured, thereby increasing predictive strength.

69 Machine learning algorithms require large datasets for the training and testing process. Traditionally, DMS measurements were based on time-consuming ship-board analysis of discrete samples, resulting in sparse data 70 71 coverage over much of the oceans. More recently, the development of several automated DMS measurement 72 systems (Royer et al., 2014; Saltzman et al., 2009; Tortell, 2005a) has provided marine DMS observations at a 73 significantly higher resolution, yielding greater spatial and temporal data coverage. These new datasets potentially 74 enable new insights into small-scale and regional patterns in oceanic DMS distributions, as well as the 75 characterization of oceanic DMS 'hot-spots'. The northeast subarctic Pacific (NESAP) is a region of notably high 76 DMS concentrations (Lana et al., 2011), with localized DMS accumulation in both highly productive coastal 77 upwelling regimes, and off-shore, iron-limited waters ((Herr et al., 2019; Asher et al., 2017). Several factors have been proposed to account for the elevated DMS production in the NESAP, including increased primary productivity 78 79 driven by nutrient entrainment and upwelling along coastal fronts (Asher et al., 2017), a dominance of high-DMSP 80 producing prymnesiophytes and dinoflagellates in offshore waters, elevated microbial degradation of DMSP to 81 DMS (Steiner et al., 2012; Rover et al., 2010), and the stimulation of DMS production in response to oxidative 82 stress in low iron waters (Sunda et al., 2002; Herr et al., 2020). Although multiple studies have examined empirical 83 relationships between DMS and various oceanographic factors in the NESAP (Watanabe et al., 2007; Herr et al., 84 2019: Asher et al., 2017, 2011), these have all reported low predictive skill based on simple linear correlation 85 approaches. To date, machine-learning approaches have not been applied to describe DMS distributions specifically 86 in this region.

87 Here, we present an approach to modelling summertime NESAP DMS concentrations and sea-air fluxes 88 using ensemble random forest regression (RFR) and artificial neural network (ANN) machine-learning algorithms. 89 Our statistical models leverage field observations of DMS collected across the NESAP between 1997 to 2017 to 90 generate a summertime DMS climatology mapped at a higher spatial resolution than previous efforts (Simó and 91 Dachs, 2002; Vallina and Simó, 2007; Galí et al., 2018; Watanabe et al., 2007; Humphries et al., 2012; Wang et 92 al., 2020). This new modelling approach represents a significant improvement over previous methods, and predicts 93 regional DMS distributions that are coherent with underlying patterns of oceanographic variability. Most notably, 94 the modelled DMS concentrations and sea-air fluxes can be explained, to a large extent, by regional and mesoscale 95 patterns in nutrient supply and physical mixing dynamics. Based on the output of our models, we present 96 summertime sea-air flux estimates in close agreement with previous studies (Herr et al., 2019; Lana et al., 2011), 97 further highlighting the importance of the NESAP as a globally-significant sulfur source to the atmosphere.

98 2 Methods

99 2.1 Data

A combination of data sources was used in training our machine-learning models to build a summertime DMS climatology. For this study, we restricted DMS measurements to the months of June, July and August between 102 1997 to 2017 in the NESAP (43-60°N, 147-122°W). A total of 26,201 data points were obtained from the NOAA 103 PMEL repository (<u>https://saga.pmel.noaa.gov/dms/;</u> last accessed: February 3, 2021), including measurements 104 derived from purge and trap gas chromatography and membrane inlet mass spectrometry. The DMS data were 105 binned to a monthly resolution, regardless of year, and averaged into 0.25 x 0.25° grid cells.

106 Predictor data used to build our machine-learning models included the following variables derived from 107 the NASA Aqua MODIS satellite at level L3 monthly 0.042° resolution (R2018.0): sea surface temperature (SST), 108 the ratio of normalized fluorescence line height to chlorophyll a (nFLH:Chl-a), instantaneous and daily observed 109 photosynthetically active radiation (iPAR and PAR, respectively), particulate inorganic carbon (PIC), the 110 absorption of gelbstof and detritus at 433 nm ($a_{cdm}(443)$), and diffuse attenuation coefficients at 490nm (K_d). Satellite-based PIC is considered as a proxy for the abundance of coccolithophores and other calcified 111 112 phytoplankton (Franklin et al., 2010), whereas the $a_{cdm}(443)$ product is considered a proxy for chromophoric dissolved organic matter (CDOM) (Nelson & Siegel, 2013), which is thought to be an important photosensitizer of 113 114 DMS (see Sect. 4.1). For observations prior to 2004, data were from either SeaWiFS (0.083° resolution) or Terra 115 MODIS (0.042° resolution) when SeaWiFS data were unavailable (e.g. nFLH and iPAR). As described below, K_d 116 and PIC were later excluded from the final models (see Sect. 2.6), as they didn't improve predictive skill.

117 The following predictor variables were also used: 6-day averaged sea surface height anomalies (SSHA) 118 derived from the TOPEX/Poseidon satellites at 0.17° resolution; Level L4 ESA Sentinal-3 Copernicus monthly-119 averaged 0.25° wind speeds; net primary productivity (NPP) from the Vertically-Generalized Production Model (VGPM; Behrenfeld & Falkowski, 1997) at monthly 0.25° resolution; sea surface nitrate from the 2018 World 120 121 Ocean Atlas at monthly 1° resolution (Garcia et al., 2019); and mixed-layer depth (MLD) and sea surface salinity 122 (SSS) from the MIMOC climatology at 0.5° resolution (Schmidtko et al., 2013). Except for MIMOC data, all 123 predictors were restricted in time to the corresponding years of DMS sampling (1997 to 2017). Net community 124 productivity (NCP) was estimated from the algorithm of Li & Cassar, (2016; using NPP and SST). As with DMS 125 observations, predictor data were interpolated to a 0.25 x 0.25° average monthly resolution using linear radial basis interpolation functions. Interpolation was constrained to the oceanic region by masking out land pixels using 126 127 ETOPO2 bathymetric (0.033° resolution) binned at 0.25 x 0.25° resolution. We note that each of these data sources 128 are likely to have inherent uncertainties associated with either their collection or processing. Data sources can be

129 found in Table 1.

130

131 Table 1. Data sources and spatial and temporal resolution of predictor variables used to develop the RFR and ANN algorithms. Data 132 processing levels are indicated where relevant. All variables were used as predictors (excluding bathymetry) and post-processed to 133 monthly-averaged, 0.25° resolution (see sections 2.1-2.2).

Variable	Spatial Resolution (°)	Temporal Resolution	Source	Level
Sea Surface Temperature (SST)	0.042	6-Day Average	SeaWiFS/AquaTERRA (1997-2003) or AquaMODIS(2004-2017): https://oceancolor.gsfc.nasa.gov/l3/	3
Chlorophyll-Normalized Fluorescence (nFLH:Chl- a)	0.042	Monthly	SeaWiFS/AquaTERRA (1997-2003) or AquaMODIS (2004-2017): https://oceancolor.gsfc.nasa.gov/l3/	3
Instantaneous Photosynthetically Active Radiation (iPAR)	0.042	Monthly	SeaWiFS/AquaTERRA (1997-2003) or AquaMODIS (2004-2017): https://oceancolor.gsfc.nasa.gov/l3/	3
Daily Photosynthetically Active Radiation (PAR)	0.042	Monthly	SeaWiFS/AquaTERRA (1997-2003) or AquaMODIS (2004-2017): https://oceancolor.gsfc.nasa.gov/l3/	3
Particulate Inorganic Carbon (Calcite; PIC)	0.042	Monthly	SeaWiFS/AquaTERRA (1997-2003) or AquaMODIS (2004-2017): https://oceancolor.gsfc.nasa.gov/l3/	3
Absorption of Gelbstof and Detritus at 433 nm (a _{cdm} (443))	0.042	Monthly	SeaWiFS/AquaTERRA (1997-2003) or AquaMODIS (2004-2017): https://oceancolor.gsfc.nasa.gov/l3/	3
Diffuse Attenuation Coefficients at 490 nm (K _d)	0.042	Monthly	SeaWiFS/AquaTERRA (1997-2003) or AquaMODIS (2004-2017): https://oceancolor.gsfc.nasa.gov/l3/	3
Sea Surface Height Anomalies (SSHA)	0.17	Monthly	TOPEX/Poseidon: <u>https://podaac.jpl.nasa.gov/dataset/SE</u> <u>A_SURFACE_HEIGHT_ALT_GRID</u> <u>S_L4_2SATS_5DAY_6THDEG_V_J</u> <u>PL1812</u>	4
Monthly Wind Speeds	0.25	Monthly	ESA Sentinal-3 Copernicus: https://resources.marine.copernicus.eu/ ?option=com_csw&view=details&pro duct_id=WIND_GLO_PHY_CLIMAT E_L4_REP_012_003	N/A

Net Primary Productivity (NPP)	0.25	Monthly	Vertically-Generalized Production Model (VGPM): <u>http://www.science.oregonstate.edu/oc</u> <u>ean.productivity/</u>	N/A
Sea Surface Nitrate (SSN)	1	Monthly	World Ocean Atlas 2018 (WO18): https://www.ncei.noaa.gov/access/worl d-ocean-atlas-2018/	N/A
Mixed Layer Depth (MLD)	0.5	Monthly	MIMOC Climatology: https://www.pmel.noaa.gov/mimoc/	N/A
Sea Surface Salinity (SSS)	0.5	Monthly	MIMOC Climatology: https://www.pmel.noaa.gov/mimoc/	N/A
Bathymetry	0.033	N/A	ETOPO2: https://rda.ucar.edu/datasets/ds759.3/	N/A

136 2.2 Machine-learning models

137 We compared the performance of random forest regression (RFR) and artificial neural network (ANN) 138 models at the regional scale. The RFR algorithm is built upon decision tree models, which operate by iteratively 139 generating decision rule nodes that dictate which branch the tree will progress through in the next iteration. The 140 RFR model builds an ensemble, or "forest", of these trees, where each tree is trained on a bootstrapped (*i.e.* 141 randomly subsampled) set of predictors, and the resulting predictions are averaged among the trees to reduce 142 overfitting to noise (Brieman, 2001). In contrast, the ANN model is built as a fully connected network of nodes, or 143 "neurons", in which each neuron consists of an activation function and is connected to other neurons by iteratively-144 determined weights (Gardner and Dorling, 1998). Both algorithms are advantageous because they make no prior 145 assumptions on the data distributions and can fit non-linear data (Brieman, 2001; Gardner and Dorling, 1998).

146 Both our ANN and RFR models followed a similar design to Weber et al. (2019). Our ANNs were built 147 using a feed-forward framework consisting of a single input node, two hidden layers each consisting of 30 neurons 148 (using a sigmoidal activation function), and a single output layer (using a linear activation function). A Bayesian 149 L2 (Ridge) regularization parameter was tuned to minimize overfitting and the L-BFGS algorithm was used to 150 solve for weights (Byrd et al., 1995). Each individual decision tree within the RFR was trained using the standard 151 CART algorithm (Brieman, 2001) and constrained to a max depth of 25 decision splits, the simplest configuration determined to perform well and minimize overfitting. These models were built using the Scikit-Learn (v0.24.2) 152 153 implementation of the ANN ("MLPRegressor") and RFR ("RandomForestRegressor") algorithms in Python 3.8 154 (see Code Availability).

In both cases, the models were built as an ensemble of either 1000 individual decision trees or individual networks to minimize bias in predictions. The input data were randomly divided for use in model training (80%) and external testing (20%). Although RFR is not sensitive to large differences in predictor variance, predictor data were standardized in both models by normalization to their respective mean and standard deviation. Additionally, we applied an inverse hyperbolic sine (IHS) transformation to the DMS data prior to training (Weber et al., 2019). Testing results indicated that IHS yielded slightly better performance than the more traditional logarithmic transformations for both of our models.

162 2.3 Sea-to-air fluxes

163 Sea-air DMS fluxes (F_{DMS} , µmol m⁻² d⁻¹) were calculated from the monthly-averaged observed and 164 modelled DMS values for June, July and August. F_{DMS} was calculated using the gas transfer velocity (k, cm hr⁻¹) 165 following the modified approach of Webb et al. (2019):

166
$$F_{DMS} = k(DMS)(0.24)$$
 (1)

where the factor of 0.24 converts to the values to daily fluxes. The gas transfer velocity has typically been calculated using a non-linear parameterization (Nightingale et al., 2000), but recent work has suggested a linear parameterization is more appropriate for DMS (Bell et al., 2013; Blomquist et al., 2017; Zavarsky et al., 2018). Since satellite-derived predictors are used to build our models, we calculated the gas transfer velocity using the linear Goddijn-Murphy et al. (2012) k parameterization, which is both derived from satellite altimeter data and normalized to a Schmidt number of 660:

173
$$k_{w,660} = 2.1U_{10} - 2.8$$
 (2)

174 Where U_{10} is the wind speed (m s⁻¹) at 10 m above sea surface.

175 Regional summertime fluxes (\bar{F}_{DMS} , Tg) were calculated as the average (±SD) quantity of DMS-sulfur 176 emitted over 92 days (June, July and August) through the area of the mapped study region (1.28x10⁷ km² or 85.0% 177 of the total bounded area).

178 2.4 Comparison against existing algorithms

Simple linear regression (LR) and multiple linear regression (MLR) models were built for comparison against the machine-learning algorithms. We also tested the performance of our RFR and ANN models against the published algorithms of Simó & Dachs (2002), Watanabe et al. (2007), Vallina & Simó, (2007), and Galí et al. (2018) (hereafter referred to as SD02, W07, VS07, and G18, respectively). Solar radiation dose, SRD, used in the VS07 algorithm was calculated using MLD as described by Vallina & Simó (2007):

184
$$SRD = \frac{PAR}{K_d \times MLD} \times (1 - e^{-K_d \times MLD})$$
(4)

Each of the four algorithms was assessed using both their original coefficients and coefficients tuned to our NESAP dataset using nonlinear least-squares optimization at both 0.25° and 1° spatial resolution (Table 2). In each case, the algorithms were run using the same monthly-averaged predictors used to develop the RFR and ANN ensembles (see Sec. 2.1). Predictors were spatially matched to either the full DMS dataset (*i.e.* all monthly averaged DMS observations) or to only the Testing partitioned dataset (see Sec. 2.2) for direct comparison with the RFR and ANN ensemble performance (Fig. 2, Table 2).

191 2.5 Controls on DMS variability

Principal component analysis (PCA) was applied to assess the relationships between DMS and the nine predictors used to build the RFR and ANN ensembles. Additionally, non-parametric Spearman rank correlations were calculated between each variable and both the modelled and observed DMS concentrations. Correlation analysis was also extended to assess the role of taxonomy on predicted DMS concentrations, using the outputs of a chlorophyll-a based taxonomic algorithm by Hirata et al. (2011) with NESAP-tuned coefficients (Zeng et al., 2018).

198 2.6 Sensitivity Tests and Predictor Selection

To inform our selection of grid size, we assessed the performance of both the RFR and ANN models using grid cells ranging from 0.25 to 5° (Fig. 1). From this analysis, we found that model accuracy was highest at 0.25° resolution (see Sect. 3.1). Smaller grid sizes would presumably further improve model accuracy, but at a significantly higher computational cost.

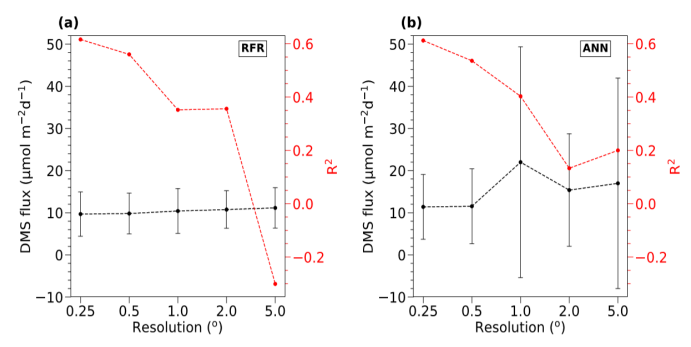
We also tested the influence of other biological predictor variables on the performance of the RFR and ANN models, using either NCP, NPP, Chl-a, or PIC. These sensitivity tests indicated no significant difference between the various biological predictor variables, although accuracy was slightly reduced when PIC was used. We therefore selected NCP as the biological predictor variable within our model framework. We also removed K_d as a predictor variable after further sensitivity testing indicated that its exclusion slightly improved results.

The inclusion of nFLH:Chl-a represents a proxy for iron limitation (see Sect. 4.1). However, fluorescence yields corrected for non-photochemical quenching (NPQ) have been suggested to yield a better iron limitation proxy than nFLH:Chl-a (Behrenfeld et al., 2009). We therefore calculated NPQ-corrected fluorescence yields (φ_f) by:

212
$$\varphi_f = \frac{nFLH}{Chl - a \times \alpha \times S} \times \frac{iPAR}{iPAR}$$
(5)

where $\alpha = 0.0147 \times Chl - a^{-0.316}$ and S = 100 mW cm⁻² µm⁻¹ sr¹ m as described by Behrenfeld et al. (2009). Our tests indicated nFLH:Chl-a yielded slightly improved performance overall, whereas φ_f decreased both models' performance. We therefore retained nFLH:Chl-a and excluded φ_f in our final model design.

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Fig. 1. Sensitivity of RFR and ANN models to grid size resolution. DMS fluxes (black) and R² values (red) derived from sensitivity tests of (a) RFR and (b) ANN models to pixels resolutions of 0.25-5°. The negative R² values observed at the lowest resolution (largest grid cells) indicate that the predicted values explain less variance than the overall mean of the dataset.

222 **3 Results**

223 3.1 Model evaluation

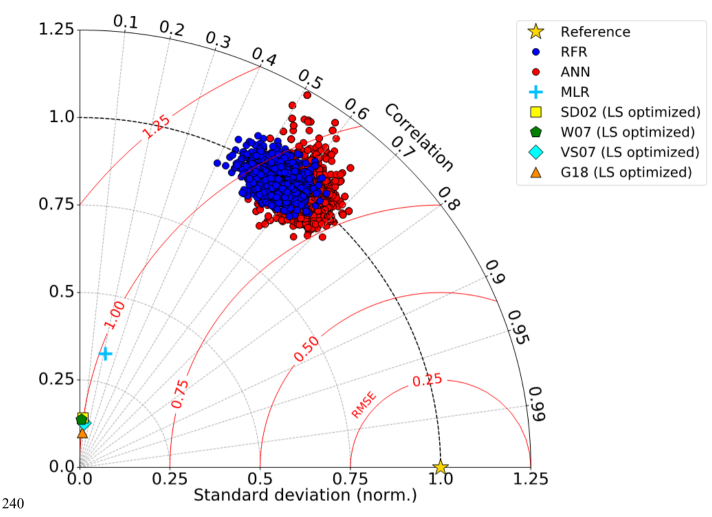
224 To benchmark the performance of our RFR and ANN models, we first evaluated the predictive skill of four 225 existing empirical DMS algorithms (SD02, W07, VS07, & G18), in addition to simple and multiple linear 226 regression models. Previous studies have demonstrated that these empirical algorithms show strong predictive skill 227 (R²=0.53-0.84) over large scales and in some oceanic regions (Simó and Dachs, 2002; Galí et al., 2018; Watanabe 228 et al., 2007), but significantly poorer performance in the NESAP (Herr et al., 2019). Consistent with these results, 229 we found that the SD02, W07, VS07, and G18 did not accurately predict NESAP DMS distributions, even with regionally tuned coefficients improving performance (Fig. 2, R²=0-0.01 at 0.25x0.25°; Table 2, r=-0.15-0.36). We 230 231 also found that simple and multiple linear regressions performed poorly, yielding virtually no explanatory power for surface water DMS distributions in the NESAP ($R^2=0-0.05$; Fig. 2, 3). 232

233

Table 2. Performance of statistical DMS algorithms on NESAP DMS observations binned to monthly 1° and 0.25° resolution. Pearson correlation coefficients (r) and root mean square error (nM) are obtained from the

SD02, VS07, W07 and G18 algorithms (see 2.4) using either their original published coefficients or
 coefficients derived from non-linear least squares optimization. Algorithm performance is evaluated using
 either the full monthly-binned observational dataset or using the Testing partitioned dataset (see Sec. 2.2).

	SD02		VS07		W07		G18	
	Original	Optimized	Original	Optimized	Original	Optimized	Original	Optimized
1° All data	r = -0.09 RMSE = 18.03	r = 0.17 RMSE = 4.82	r = -0.03 RMSE = 6.67	r = 0.03 RMSE = 4.96	r = -0.10 RMSE = 11.74	r = 0.07 RMSE = 4.83	r = 0.02 RMSE = 6.77	r = 0.16 RMSE = 4.84
1° Testing dataset	r = -0.22 RMSE = 19.09	r = 0.36 RMSE = 3.34	r = 0.11 RMSE = 5.36	r = 0.20 RMSE = 3.47	r = -0.03 RMSE = 10.46	r = 0.02 RMSE = 3.47	r = -0.15 RMSE = 6.19	r = 0.30 RMSE = 3.40
0.25° All data	r = -0.05 RMSE = 11.02	r = 0.12 RMSE = 7.84	r = -0.09 RMSE = 9.57	r = 0.11 RMSE = 7.88	r = -0.09 RMSE = 13.02	r = 0.04 RMSE = 7.80	r = 0.06 RMSE = 8.42	r = 0.09 RMSE = 7.88
0.25° Testing dataset	r = -0.03 RMSE = 9.79	r = 0.07 RMSE = 6.79	r = -0.09 RMSE = 8.60	r = 0.10 RMSE = 6.79	r = -0.06 RMSE = 12.02	r = 0.04 RMSE = 6.78	r = 0.04 RMSE = 7.47	r = 0.08 RMSE = 6.80



241 Fig. 2. Taylor Diagram showing comparative performance metrics of each individual Random Forest Regression (RFR) 242 and Artificial Neural Network (ANN) model (1000-model ensembles) against multiple linear regression (MLR) and 243 other statistical DMS models (See sections 2.1 and 2.4). The Pearson correlation coefficients ("Correlation"; outer 244 radius), root mean squared error ("RMSE"; red radial contours), and standard deviations (SDs; grey radial contours 245 from origin) are all computed with respect to the observed DMS samples after inverse hyperbolic sine (IHS) 246 transformation. The reference of a perfect model fit is shown with a gold star. SDs of the model outputs are normalized 247 to the SDs of the DMS observations. RMSE represents a normalized trigonometric derivation from both the correlation 248 coefficients and normalized SDs. Performance of the SDO2, W07, VS07, and G18 algorithms reported here are 249 calculated using regionally tuned coefficients to the NESAP derived from non-linear least-squares optimization (see 250 section 2.4).

Relative to other published modelling approaches, both the RFR and ANN models dramatically improved the representation of NESAP DMS variability, achieving significantly higher predictive accuracy (Fig. 2, 3). The collective ensembles of both the RFR and ANN models yielded strong performance, explaining up to 62% of the observed DMS variability ($R^2=0.61-0.62$; Fig. 3). For individual models within the ensembles, the ANN method provided slightly better results ($R^2=0.16-0.50$), compared to the individual RFR models ($R^2=0.16-0.43$). However, predicted DMS concentrations and sea-air fluxes derived from the ANN ensembles were more sensitive to the spatial resolution used, although the predictive accuracy of both models degraded significantly with coarser resolutions (Fig. 1).

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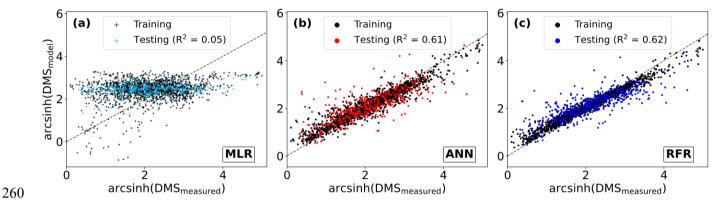


Fig. 3. Performance of three modelling approaches in predicting observed DMS distributions; (A) multiple linear regression (MLR) (B) ensemble of Artificial Neural Networks (ANN) and (C) ensemble of Random Forest Regression (RFR). For consistency, all predictions are partitioned by the Training and Testing datasets used to build the ensembles (see section 2.2). Model performance (R²) is computed only for the Testing dataset predictions. The dashed line demonstrates a 1:1 relationship. Modelled DMS concentrations depicted range from 0.4-84.3 (RFR, nM) and 0.3-74.6 (ANN, nM).

267 3.2 DMS distributions and sea-air fluxes

268 In both the RFR and ANN methods, the predicted spatial distribution of DMS was generally consistent with 269 observations (Fig. 4a,c,d). The average model derived DMS concentrations was 4.0 ± 2.1 nM and 4.7 ± 3.0 nM 270 (mean \pm SD) for the RFR and ANN ensemble models, respectively, with a similar range from 0.3 to 84.3 nM. In 271 both models, the highest DMS concentrations were largely constrained to coastlines and within the Alaska Gyre 272 adjacent to the Aleutian Islands (Fig. 4b-c, 8C). The greatest discrepancy between DMS concentrations from the two models was observed in these regional 'hotspots', where the ANN models emphasize high DMS within the 273 274 Alaska gyre, while the RFR models emphasize elevated coastal DMS concentrations (Fig. 4b). On average, the models deviated from each other by 0.49 nM, with the greatest offsets observed in an area of particularly sparse 275 276 DMS observations in the Alaska Gyre (Fig. 4a,b). Future observational data in this region should help improve 277 model agreement.

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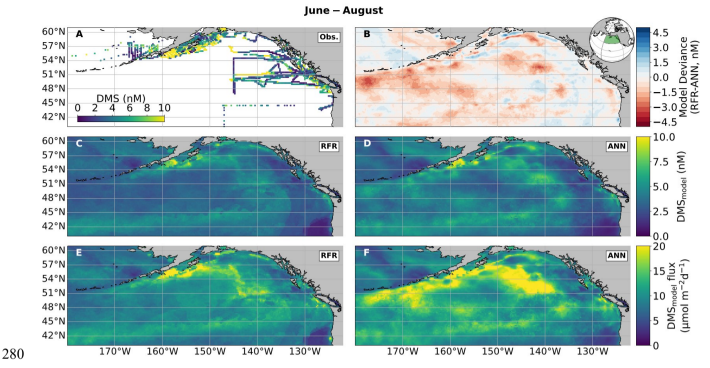


Fig. 4. Predicted maps of sea surface DMS concentrations and sea-air fluxes. (a) Ship-based observations of mean summertime (June-August) DMS concentrations used to construct the predictive models. (b) Differences between the (c) Random Forest Regression (RFR) and (d) Artificial Neural Network (ANN) ensemble predicted DMS concentrations. (e,f) DMS sea-air fluxes derived from the predicted DMS concentrations. Colormap ranges are restricted to illustrate trends, with <1% of DMS data exceeding the colorbar limits. The inset map in (b) shows the NESAP study region as a shaded green patch in a global orthographic projection.

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288 Sea-air DMS fluxes (Fig. 4e,f) derived from ANN predictions were 18% higher, on average, than RFR 289 predictions, largely due to higher predicted values in the Alaska Gyre (Fig. 4d-e, Table 3). The distribution of ANN 290 sea-air fluxes was also closer to ship-based observations (Fig. 5). Predicted regional fluxes ranged from 0.8 to 167 291 umol m^{-2} d⁻¹ between the two models (Fig. 4e,f. 5), with the highest predicted DMS emissions in August, when 292 derived sea-air fluxes were approximately 1.6 to 2-fold greater than in June and July (Table 3). Our models yielded 293 a summertime integrated sea-air flux of 1.16±1.22 Tg DMS-derived sulfur, which is consistent with the Lana et al. 294 (2011) climatological estimate of 1.64 ± 0.51 Tg (Table 3). 295 Table 3. Monthly and mean summertime NESAP sea-air DMS fluxes. Total cumulative fluxes of DMS-derived sulfur (Tg,

296 mean \pm SD) are calculated from the Random Forest Regression (RFR) and Artificial Neural Network (ANN) model predictions

297 (based on an ensemble of 2000 models). Total cumulative NESAP sea-air flux derived from the Lana et al. (2011) climatology

298 is shown for comparative purposes.

Summertime Sulfur Emissions

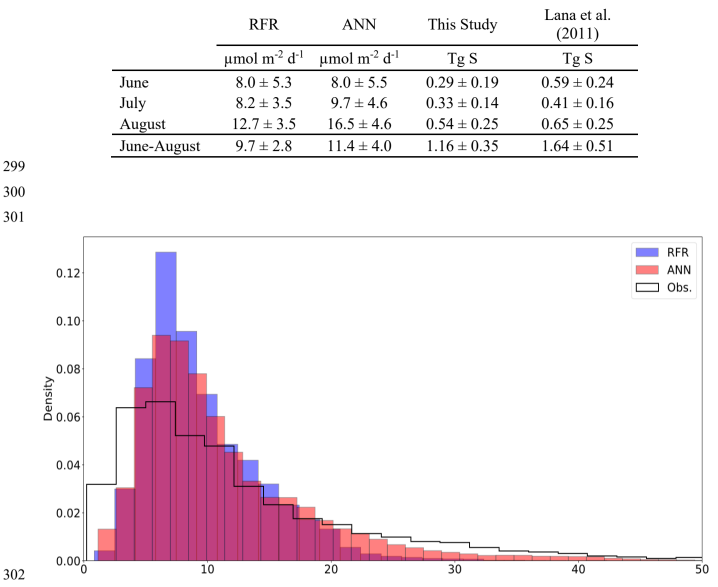
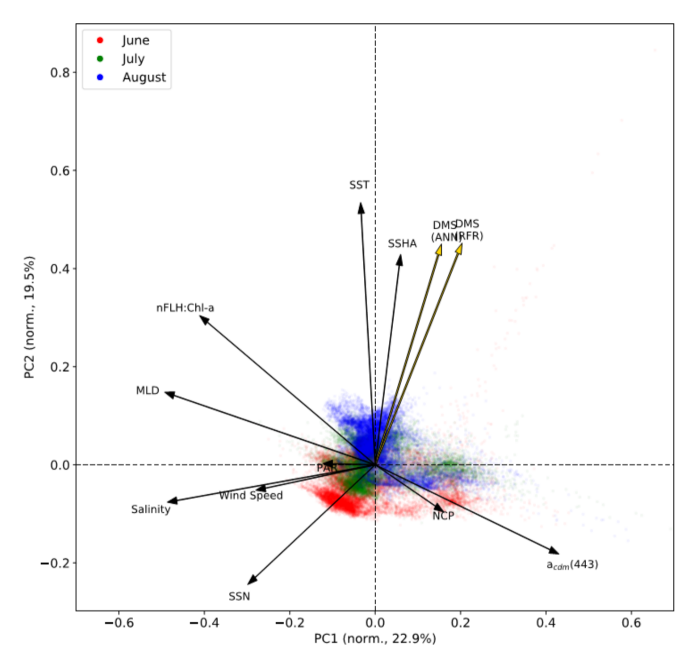


Fig. 5. Histograms of DMS sea-air flux distributions derived from the 1000-model ensemble random forest regression (RFR) and artificial neural network (ANN) predictions as well as cruise observations (Obs.). The sample sizes of both models are equivalent (n= 49,632) and are significantly higher than the observational dataset (n=2063). Note that the distribution is restricted to show trends, with a maximum flux of 238 nM (Obs.). The upper tail (>50 nM) consists of only 2.9% (Obs.) and <0.1% (both RFR and ANN) of the values. Note that the ANN better predicts the upper tail of DMS observations greater than 20 nM.

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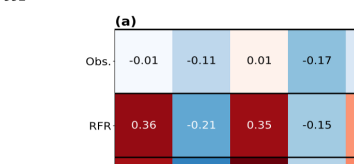
309 3.4 Drivers of DMS variability

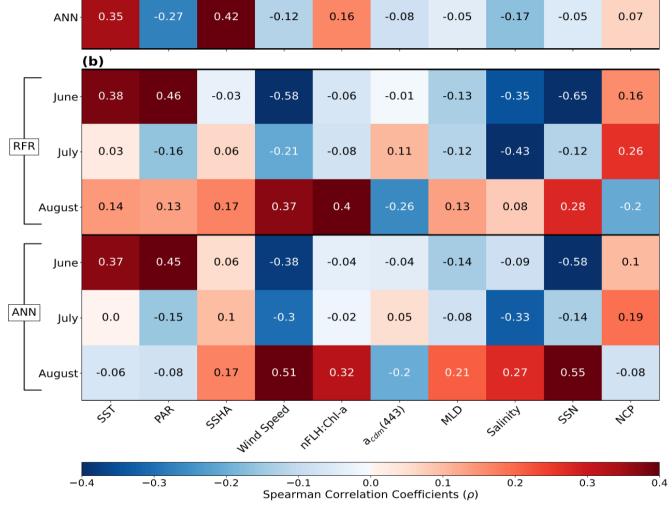
310 In addition to modelling the spatial and temporal distribution of surface water DMS in the NESAP, we 311 examined the influence of different oceanographic variables as model predictors. As expected based on previous 312 work (Herr et al., 2019), no single predictor was found to exert a dominant control on modelled DMS distributions 313 from either the RFR or ANN models (Fig. 6, 7). Rather, the relationship between DMS and other oceanographic 314 variables exhibited significant region-specific patterns. One of the most compelling regional signatures was the 315 apparent relationship between DMS and SSHA. In both models, we found significant positive correlations between 316 DMS and SSHA (ρ =0.35, 0.42 for RFR and ANN, respectively) across the full spatial domain, with a particularly 317 notable relationship along the northern Alaskan coastline (Fig. 8, 9). Here, strong winds (Fig. 9j-1), coupled with 318 the northeastern Alaska current flow, produce two characteristic oceanographic features in the NESAP: strong, 319 semi-permanent mesoscale eddies collectively referred to as the Haida, Sitka and Yakutat eddies (Fig. 8a), and the 320 formation of the high nutrient, low chlorophyll (HNLC) Alaska Gyre (Fig. 8c; Okkonen et al., 2001; Whitney et 321 al., 2005). Both the monthly (Fig. 9a-i) and summertime-averaged (Fig. 8a,b) RFR and ANN-derived DMS 322 concentrations are low where these downwelling eddies form. In contrast, elevated DMS concentrations were 323 associated with the negative SSHA coastal upwelling areas (Fig. 8a,b), where phytoplankton productivity is 324 stimulated by nutrient inputs into the mixed layer.



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Fig. 6. Principal Component Analysis (PCA) showing the relationships between variables used to construct the predictive algorithms. Eigenvectors (arrows) are superimposed over the principal components (PCs; data points) for the first two significant modes obtained from PCA. PCs are normalized and clustered by month (June-August, see legend for colors), while the eigenvectors are grouped by ensemble model predictions (gold) and nine predictor variables (black). The percentage of variance explained by each mode is indicated along the axes.





-0.06

0.15

0.03

-0.04

-0.04

-0.09

-0.08

-0.26

0.02

-0.15

0.08

0.08

333

Fig. 7. Heatmap of Spearman rank correlations (ρ). (a) Correlations of pooled data (June-August) for DMS
observations (Obs.), RFR and ANN predictions per variable. (b) Correlations per month for the RFR and ANN DMS
predictions. All model correlations are computed on the 1000-model ensembles.

337 Modelled DMS concentrations also significantly correlated with hydrographic frontal patterns. We found 338 significant correlations between DMS and SST ($\rho=0.36$, 0.35 for RFR and ANN, respectively) which suggested 339 the central Alaska Gyre and offshore of Vancouver Island are areas of elevated DMS variability (Fig. 8b). Both 340 models predict high DMS levels in the northern frontal zone of the gyre (140°W-145°W) between the 10.5 and 12°C isotherms and the southern frontal zone between (42°N-45°N) between the 13.5 and 15°C isotherms (Fig. 341 342 8b,c). By comparison, our models suggest that DMS concentrations are predominantly low in relation to high sea 343 surface nitrate (SSN) concentrations within the HNLC gyre (Fig. 8, 9). As discussed below, the relationship 344 between DMS and macronutrient concentrations in the HNLC waters of the central Gulf of Alaska could indicate 345 an important role for iron limitation as a controlling factor in the DMS cycle. The presence of elevated summer nutrients in offshore waters is taken as a proxy for iron limitation, which increases over the course of the summer 346 347 growing season.

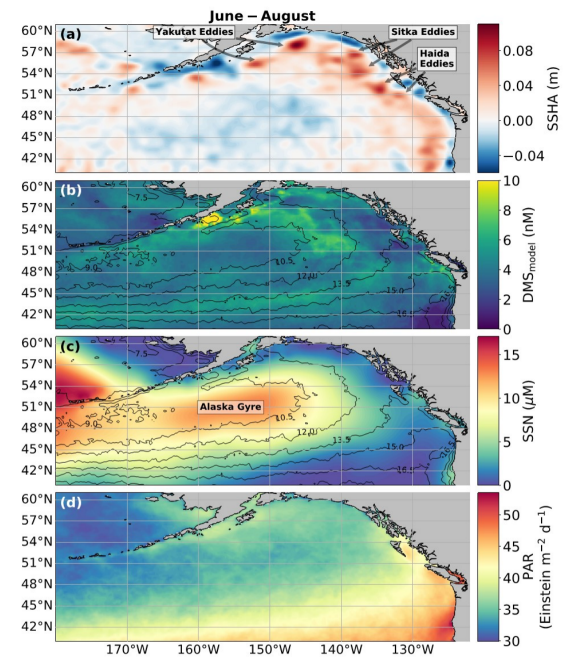




Fig. 8. Physical drivers of summertime (June-August) NESAP DMS distributions. (a) Sea surface height anomalies (SSHA), (b) predicted DMS concentrations derived from the mean of all 2000 RFR and ANN machine learning models, (c) sea surface nitrate (SSN) and (d) photosynthetically active radiation (PAR). Contours in (b,c) show sea surface temperature (SST) isotherms. Coherent features of elevated sea-surface height indicate the presence of mesoscale eddies, whereas nearshore low SSHAs features reveal areas of upwelling. Colormaps ranges are restricted to illustrate trends with <1% of data exceeding the colorbar limits.</p>

355 Other variables appear to exhibit a more localized or minimal influence on DMS cycling. For instance, 356 both NCP and DMS are elevated in productive nearshore waters, but NCP generally correlates weakly with both 357 RFR- and ANN-derived DMS concentrations ($\rho=0.08$, 0.07 for RFR and ANN, respectively). It should be noted, 358 however, the empirically-derived NCP estimates may carry more uncertainty than other predictors obtained from 359 direct satellite observations (Li and Cassar, 2016). Similarly to NCP, modelled phytoplankton taxonomic 360 composition (Hirata et al., 2011; Zeng et al., 2018) was not significantly correlated with predicted DMS 361 concentrations (ρ <0.1). Although strong, persistent winds appear to sustain low DMS concentrations off the coast 362 of Oregon and Vancouver Island (Fig. 9), wind speeds only weakly correlate with DMS overall for the region (p=-363 0.15 and -0.12 for RFR and ANN, respectively). Additionally, high PAR in these areas correspond with low DMS 364 concentrations (Fig. 6d) and there is an overall negative correlation between PAR and DMS for the region (Fig. 6, 365 7; ρ =-0.21 and -0.27 for RFR and ANN, respectively). Finally, despite hypothesized links between DMS cycling 366 and iron limitation in the NESAP (Levasseur et al., 2006; Merzouk et al., 2006; Rover et al., 2010), nFLH:Chl-a 367 ratios (taken as a proxy for phytoplankton iron stress; Behrenfeld et al., 2009; Westberry et al., 2013) did not exhibit 368 any coherent spatial patterns, and only weakly correlated to our modelled DMS concentrations ($\rho=0.15$ and $\rho=0.16$ 369 for RFR and ANN, respectively).

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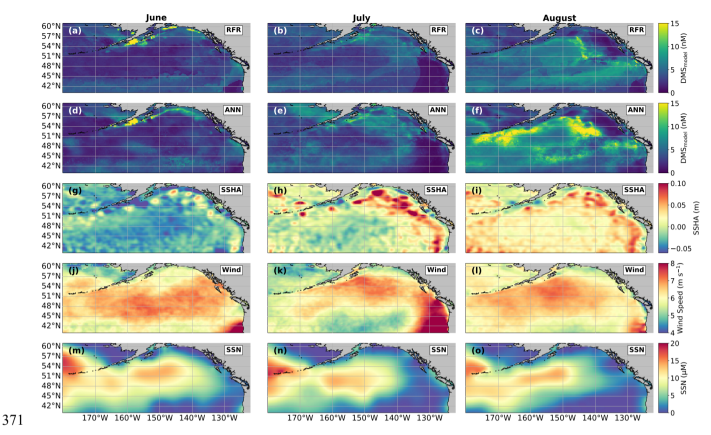


Fig. 9. Predicted spatial and temporal (June-August) DMS distribution in relation to underlying oceanographic variables. DMS concentrations predicted from (a-c) the Random Forest Regression (RFR) and (d-f) the Artificial Neural Network (ANN) ensemble models are mapped alongside the monthly-averaged (g-i) sea surface height anomalies (SSHA), (j-l) wind speeds (Wind), and (m-o) sea surface nitrate (SSN) for each month. Colormap ranges are restricted to illustrate trends, with at most 1.5% of the data beyond the colorbar limits.

377 4 Discussion

378 The relative sparsity of DMS data in many oceanic regions and the complexity of DMS cycling have limited 379 previous attempts to model oceanic distributions of this compound (Simó and Dachs, 2002; Vallina and Simó, 380 2007; Galí et al., 2018; Watanabe et al., 2007; Herr et al., 2019). Taking advantage of expanding data resources, 381 we employed a new approach to statistically describe DMS distributions in the NESAP. Our results show that both 382 our RFR and ANN models substantially improved predictive strength over traditional empirical approaches (Fig. 383 2, 3), while identifying several key DMS relationships and regional patterns across the NESAP (Fig. 8, 9). Although 384 our statistical approach does not directly elucidate the underlying mechanisms driving these relationships, and not 385 all variability in predictors may be captured at the single spatial scale used here, we can nonetheless make some 386 reasonable inductive inferences. These inferences are discussed below, along with the implications of the improved

387 predictive performance observed here.

388 4.1 Relationships with other oceanographic variables

389 Among the more prominent spatial relationships we observed was the coherence between predicted DMS 390 concentrations and SST, and the negative correlation between predicted DMS concentrations and sea surface nitrate (SSN) within and surrounding the Alaska Gyre (Fig. 6-9). Notably, regional SSN, NCP and Chl-a distributions did 391 392 not vary appreciably inside versus outside the gyre, and these variables were poorly correlated with DMS 393 concentrations (r=-0.02, ρ =0.08 with NCP, r=0.09, ρ =-0.12 with Chl-a). This suggests that the patterns in surface 394 DMS across the Alaska Gyre were not simply driven by changes in phytoplankton biomass or productivity. The DMS-nitrate relationship may be partially explained by the so-called sulfur overflow hypothesis (Stefels, 2000), 395 396 which suggests that nutrient-limited phytoplankton increase DMSP production and its subsequent cleavage to 397 DMS, in order to regulate intracellular sulfur quotas when protein synthesis is limited (Hatton & Wilson, 2007; 398 Kinsey et al., 2016; Simó & Vila-Costa, 2006; Spiese & Tatarkov, 2014; Stefels, 2000). This mechanism may help 399 explain the higher predicted DMS concentrations at the northern extent of the Alaska Gyre, where SSN 400 concentrations begin to decrease (Fig. 6). Nutrient-dependent effects may also be important in explaining seasonal 401 variability, as the DMS-nitrate relationship becomes positive in August as phytoplankton growth becomes 402 increasingly nutrient limited (Fig. 7b).

The apparent relationship between DMS and nitrate could also result indirectly from the underlying effects of iron limitation. Excess summertime nitrate concentrations are taken as evidence for iron limitation in the NESAP (Boyd and Harrison, 1999; Boyd et al., 2004; Martin and Fitzwater, 1988; Whitney et al., 2005). Under iron-limiting conditions, DMS is thought to function, together with DMSP and DMSO, as part of an antioxidant response to oxidative stress (Sunda et al., 2002). This hypothesis suggests that iron limitation should stimulate net production of DMS and DMSP (Bucciarelli et al., 2013; Sunda et al., 2002), which is inconsistent with the overall negative dependence predicted between DMS and SSN (Fig. 8b,c).

Satellite-based, chlorophyll-normalized fluorescence has been suggested as an additional proxy for iron limitation. Low iron conditions can lead to both a reduction in photosystem I relative to photosystem II (Strzepek and Harrison, 2004), and an apparent increase in energetically-decoupled light harvesting complexes (Allen et al., 2008; Behrenfeld & Milligan, 2013), resulting in elevated fluorescence-to-chlorophyll a ratios (nFLH:Chl-a) (Westberry et al., 2013). To our knowledge, this proxy has not been widely investigated with respect to DMS cycling. In our analysis, we found that nFLH:Chl-a ratios, and the NPQ-corrected fluorescence yields (φ_f), exhibited 416 only weak positive correlations with the RFR and ANN predicted DMS concentrations (Fig. 6, 7). Moreover, 417 neither of these metrics exhibited coherent spatial patterns with predicted DMS concentrations, suggesting a limited 418 role for iron in driving spatial patterns of DMS cycling within the NESAP. However, it is important to note the 419 potential temporal mismatch between our monthly DMS predictions and these more instantaneous metrics of iron 420 limitation, which reflect short-term physiological changes (days to weeks; Behrenfeld et al., 2009; Westberry et 421 al., 2019) that depend on sporadic iron loading (e.g. aerosol deposition; Mahowald et al., 2009). Indeed, both natural 422 and artificial iron-fertilization events have thus far been detected from satellite-derived nFLH:Chl-a at daily 423 resolution (Westberry et al., 2013), in contrast to the monthly-averaged data used here. Therefore, modelling 424 frameworks utilizing shorter temporal scales may find a clearer connection between DMS cycling and iron 425 limitation using the chlorophyll-a fluorescence proxy.

426 Beyond nutrient limitation effects, ambient light fields are believed to exert significant direct and indirect 427 effects on DMS cycling (del Valle et al., 2007). At the community level, high irradiance may inhibit bacterial consumption of DMS (Slezak et al., 2001; Toole et al., 2006; Lizotte et al., 2012), while covarying changes in 428 429 mixing and high irradiance can induce transient selectivity for high-light acclimated species and influence the proportion of high DMS/P producers within assemblages (Galí et al., 2013; Vance et al., 2013). Ultraviolet 430 431 radiation has been noted to induce high DMS production and turnover through a proposed cascading oxidation 432 pathway, which acts to remove harmful reactive oxygen species (Sunda et al., 2002; Archer et al., 2010). In contrast, 433 more recent evidence has indicated the potential for elevated DMS production in the NESAP from the reduction of 434 DMSO due to light-induced oxidative stress over diurnal cycles (Herr et al., 2020). Although our modelled DMS 435 concentrations exhibited an overall negative correlation with PAR (Fig. 6, 7a), monthly correlations indicate a 436 stronger positive correlation between DMS and PAR in June, where the summer solstice drives high irradiance. In 437 contrast, July and August exhibit much weaker negative correlations as the summer bloom declines (Fig. 7b). These 438 results provide indirect evidence that light-induced oxidative stress, possibly coupled with inhibition of microbial 439 DMS consumption, may influence regional NESAP DMS distributions, particularly early in the summer.

The overall negative association of DMS and incident light (Fig. 6,7a) may also indicate a role for photolysis in DMS loss through (del Valle et al., 2007). Since DMS does not have strong light absorption properties, the presence of photosensitisers is necessary for the abiotic photooxidation of DMS (Brimblecombe and Shooter, 1986). To account for this process, our models incorporated nitrate (SSN) and a_{cdm}(443) (as a proxy for CDOM; Nelson & Siegel, 2013), both of which are thought to be dominant photosensitisers of DMS in marine systems (Taalba et al., 2013; Bouillon and Miller, 2004, 2005; Galí et al., 2016). In the NESAP, nitrate appears to exert a stronger influence than CDOM on the apparent quantum yields (AQY) of DMS (Bouillon and Miller, 2004). In support of this, our results suggest a stronger negative dependence of predicted DMS concentrations on nitrate compared to CDOM within the NESAP, particularly in June when irradiance is high (Fig. 6, 7). We acknowledge, however, that the DMS-nitrate relationship likely also reflects physiological impacts of nutrient limitation, as discussed above. Nonetheless, our results are consistent with elevated rates of DMS photo-oxidation in the nitratereplete, low iron waters of the Alaska Gyre, where photolysis may drive strong DMS oxidation and explain the low predicted DMS concentrations (Fig. 8, 9). Further *in situ* work will be required to resolve the relative contributions of these biotic and abiotic processes to DMS cycling within these areas.

454 Among all the statistical relationships we observed, perhaps the most striking was the association of DMS 455 variability with SSHA, particularly along the Alaskan coast and in relation to mesoscale eddies (Okkonen et al., 456 2001: Whitney et al., 2005: Fig. 8, 9). To our knowledge, only one other study has linked SSHA to DMS within 457 the NESAP. Herr et al., (2019) demonstrated contrasting positive and negative correlations between DMS and 458 SSHA in offshore and coastal waters, respectively, in general agreement with our results. Presently, the underlying mechanisms explaining the relationship between SSHA and DMS cycling remain unclear, yet it is likely that 459 460 physical mixing processes are important. For example, enhanced biological production is known to be stimulated by eddy re-supply of iron and macronutrients via vertical advection and diffusion (Whitney et al., 2005; Bailey et 461 462 al., 2008). These nutrient supply processes would also be expected to influence DMS cycling, as outlined above. 463 Elevated abundances of high DMS-producers within anticvclonic eddies with positive sea surface height anomalies 464 have been noted in the Sargasso Sea (Bailey et al., 2008), while eddy-induced vertical transport likely supplements 465 nearshore, current-driven upwelling that can also resupply iron into the coastal waters of the NESAP (Cullen et al., 466 2009; Freeland et al., 1984). In addition, eddy propagation can allow cross-shelf transport, distributing 467 micronutrients to offshore waters (Fiechter and Moore, 2012), potentially contributing to the apparent elevated 468 DMS concentrations in the outer Alaska gyre between the 10.5 and 12°C isotherms (Fig. 8). These mixing and 469 transport mechanisms could partially explain the influence of elevated productivity in driving increased nearshore and northern NESAP DMS concentrations (Fig. 4, 7-9), representing a novel source of DMS variability in this 470 471 region.

The taxonomic composition of plankton assemblages is also a likely source of variability influencing DMS cycling. Significant changes to DMS production and consumption rates within the NESAP are expected in response to variable microbial and phytoplankton taxonomy (Vila-Costa et al., 2006; Lidbury et al., 2016; Sheehan and Petrou, 2020). Such taxonomic variability may, in turn, reflect transient community composition shifts in response to mixing (Bailey et al., 2008), nitrate (Bouillon and Miller, 2004), and iron availability (Levasseur et al., 2006; Merzouk et al., 2006). The monthly averaging used in our data processing removes autocorrelation associated with 478 individual sampling expeditions (Wang et al., 2020), but it may preclude capturing these transient taxonomic 479 responses. For instance, coccolithophores are believed to influence DMS cycling in the NESAP (Herr et al., 2019; 480 Asher et al., 2011), yet monthly-averaged calcite distributions did not yield increased predictive strength for DMS 481 concentrations in our analysis (see Sect. 2.6). However, as satellite PIC preferentially reflects the optical signature 482 of detached coccoliths, monthly-averaged satellite PIC observations may represent the senescence of 483 coccolithophore blooms, rather than active growth phases. Additionally, applying a chlorophyll-a based taxonomic 484 algorithm (Hirata et al., 2011; Zeng et al., 2018) yielded no further explanation of the DMS variability predicted. 485 The influence of taxonomic composition thus remains cryptic within our modelling framework.

486 4.2 Implications of Improved Predictive Power

487 As noted above, both the RFR and ANN approaches demonstrate significantly improved accuracy over 488 existing models, explaining up to 62% of observed DMS variability (Fig. 2, 3). This predictive skill is somewhat 489 lower than that achieved for methane fluxes (Weber et al., 2019) and dissolved inorganic carbon dynamics (Roshan 490 and DeVries, 2017), where R^2 values ranging from 0.7 to 0.95 were obtained. Nonetheless, the dramatic accuracy 491 improvement of our algorithms over traditional methods (Fig. 2, 3) encourages the further use of these techniques 492 in modelling DMS distributions.

493 Improved predictive accuracy provides opportunities to gain insight into the mechanisms driving DMS cycling. 494 Our approach has yielded accurate DMS predictions at a 4 to 40-fold higher resolution then previous algorithms 495 (Simó and Dachs, 2002; Vallina and Simó, 2007; Galí et al., 2018; Watanabe et al., 2007), enabling the description 496 of mesoscale patterns and processes (Fig. 8). Extending these methods to sub-mesoscale resolution will enable 497 investigations into the dependence of DMS on finer-scale hydrographic processes, particularly stratification and 498 frontal dynamics, which have been increasingly linked to DMS cycling but remain unresolved mechanistically 499 (Rover et al., 2015; Asher et al., 2011). Moreover, coupling machine learning algorithms with biophysical and 500 tracer export models holds promise to resolve the contributions of eddy dynamics and upwelling intensity on DMS 501 variability, likely through nutrient availability and physiological mechanisms (Asher et al., 2011; Bailey et al., 502 2008; Cullen et al., 2009). Recent work has also developed a new database of DMS apparent quantum vields (Galí 503 et al., 2016). As the availability of these measurements increases, simultaneous mapping of both DMS quantum 504 yields and concentrations will become feasible, enabling future studies to better parse out the contribution of 505 photolysis, physical mixing, and biological drivers of DMS cycling.

Although used in a diagnostic capacity here, our statistical models also hold potential for prognostic
 applications. Frameworks utilizing shorter time scales will likely be able to detect underlying mechanisms

508 driving observed diel cycling (Galí et al., 2013; Rover et al., 2016), even if the underlying mechanisms are still 509 unresolved. We note, however, that caution will need be exercised as machine learning models have a tendency 510 to overfit noise (Weber et al., 2019; Roshan and DeVries, 2017; Wang et al., 2020), thus requiring appropriately 511 large training datasets and the use of known "future" observations to validate predictive accuracy in this context. 512 The significant variability in DMS cycling across oceanic regimes will likely also render predictions more 513 successful at regional, rather than global, scales (Galí et al., 2018; Royer et al., 2015). Nonetheless, prognostic 514 applications of these algorithms should be investigated to aid in the future development of improved mechanistic 515 models.

516 **5 Conclusions**

517 We have presented a statistical approach for modelling DMS distributions, which provides significantly 518 higher predictive skill than traditional methods (Simó and Dachs, 2002; Vallina and Simó, 2007; Galí et al., 2018; 519 Watanabe et al., 2007; Lana et al., 2011), and yields estimates of the summertime NESAP DMS sea-air fluxes to 520 1.16±1.22 Tg S in agreement with previous findings (Herr et al., 2019; Lana et al., 2011). Our results further 521 underscore the importance of the NESAP to global DMS production and motivate further observations in 522 traditionally under-sampled areas such as the Alaska Gyre and Aleutian Islands. Although we are unable to directly examine the mechanistic drivers of DMS variability, our findings suggest nutrient limitation, light-driven 523 524 processes, and eddy-induced mixing are potentially key drivers of DMS cycling in the NESAP. Future studies will 525 benefit from using such statistical algorithms, in conjunction with field-based process studies and mechanistic 526 models, to better understand the underlying dynamics and driving factors in the oceanic DMS cycle.

527 Code availability. The analysis in this study makes extensive use of the Numpy, Matplotlib, & Scikit-Learn libraries 528 codes used be in Pvthon. The custom can downloaded at 529 https://github.com/bjmcnabb/DMS Climatology/tree/main/NESAP or are available upon request from the 530 corresponding author.

531 *Data Availability.* DMS observations and predictor datasets are described in the Methods with relevant links to 532 repositories. Data from the Lana et al. (2011) climatology used for comparison in Table 3 are available via the 533 SOLAs project (retrieved from <u>www.bodc.ac.uk/solas_integration/implementation_products/group1/dms/</u>), where 534 the DMS sea-air fluxes were calculated as described in Sect. 2.3. The gridded climatologies produced from each

535algorithminthisstudycanbeobtainedat536https://github.com/bjmcnabb/DMSClimatology/tree/main/NESAP/Climatologies.at

Author Contribution. BM and PT designed the study. Model code was written and implemented by BM. BM
 prepared the manuscript with significant contributions from PT.

539 Competing Interests. The authors declare that they have no conflict of interest.

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